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Geographic variation in the time-frequency characteristics of high-frequency whistles produced by killer whales (*Orcinus orca*)

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Abstract

Investigating intra-specific variation in acoustic signals can indicate the extent of isolation and divergence between populations and adaptations to local environments. Here we analyze the variation in killer whale high-frequency (>17 kHz) whistles recorded off Norway, Iceland, and the North Pacific. We used a combination of methods including multivariate comparisons of spectral and temporal parameters and categorization of contours to types. Our results show that spectral and temporal characteristics of high-frequency whistles recorded in the North Pacific show significant differences from whistles recorded in the Northeast Atlantic, being generally stereotyped, lower in frequency, and slightly longer in duration. Most high-frequency whistles from the North Pacific were downsweeps, while this was one of the least common types recorded in the Northeast Atlantic. The repertoire of whistles recorded in Norway was similar to Iceland, but whistles produced in Norway had significantly lower maximum frequency and frequency range. Most methods were able to discriminate between whistles of the North Pacific and the Northeast Atlantic, but were unable to consistently distinguish whistles from Iceland and Norway. This suggests that macro- and microgeographic differences in high-frequency whistles of killer whales may reflect historical geographic isolation between ocean basins and more recent divergence between adjacent populations.

Keywords: ultrasonic whistles, orcas, geographic variation

40 Introduction

41 When describing geographic variation in acoustic signals it is useful to divide
42 differences into macrogeographic or microgeographic variation, depending on
43 whether populations being compared are neighboring, making interbreeding and
44 social interaction a possibility, or geographically separated, and thus socially and
45 reproductively isolated (Mundinger 1982). Comparative studies of odontocete
46 whistles have shown more pronounced inter- than intra-specific variation in whistle
47 spectral parameters, which could arise if intra-specific variability is constrained to
48 occur within a species-specific framework (Steiner 1981, Ding *et al.* 1995a, Matthews
49 *et al.* 1999, Rendell *et al.* 1999, Podos *et al.* 2002). Variation in whistle structural
50 parameters has previously been described for populations of the same dolphin species
51 both at microgeographic (*e.g.*, Azevedo and Van Sluys 2005, Morisaka *et al.* 2005)
52 and macrogeographic levels (*e.g.*, Camargo *et al.* 2006, Baron *et al.* 2008). Intra-
53 specific geographic variations may occur due to genetic divergence, adaptations to
54 local environments or cultural differences (Janik 2009). A general correlation of
55 whistle variation with distance has been proposed whereby neighboring populations
56 appear to have more similar whistle structure than geographically distant populations
57 (Ding *et al.* 1995b, Azevedo and Van Sluys 2005, Rossi-Santos and Podos 2006,
58 May-Collado and Wartzok 2008), although this is not always the case (Camargo *et al.*
59 2006). Within populations, there is often variation in duration and complexity of
60 signals (*i.e.*, number of inflection points) (Steiner 1981, Azevedo and Van Sluys
61 2005, Morisaka *et al.* 2005), while the time-frequency contour of whistles often varies
62 among individuals, providing information on individual identity (*e.g.*, bottlenose
63 dolphin, *Tursiops truncatus*; Janik *et al.* 2006; common dolphin, *Delphinus delphis*:
64 Caldwell and Caldwell 1968; Pacific humpback dolphin, *Sousa chinensis*: Van Parijs

65 and Corkeron 2001). Understanding the variation in the acoustic repertoire within and
66 between different populations may shed light on the evolution, movements, and
67 cultural traditions of a species.

68 There are known differences in the structure and use of pulsed calls and whistles
69 between different killer whale populations. In the North Pacific, pods of resident
70 (fish-eating) killer whales produce unique and stable repertoires of stereotyped pulsed
71 calls (Ford 1989, 1991). These calls differ between matriline within pods (Ford 1991,
72 Miller and Bain 2000, Deecke *et al.* 2010), and to a lesser degree between individuals
73 within the same matriline (Nousek *et al.* 2006). The stability of resident pods may
74 explain why the variation in pulsed calls primarily encodes group, rather than
75 individual, identity (Tyack 1986). Killer whale whistles are generally more complex
76 and longer than other delphinid whistles (Thomsen *et al.* 2001) and some have
77 stereotyped frequency contours that are often produced in complex sequences (Riesch
78 *et al.* 2006, 2008). Resident killer whale groups in British Columbia that do not share
79 pulsed calls share stereotyped whistles, which may provide a community-level means
80 of recognition, facilitating social interactions (Riesch *et al.* 2006). Transient
81 (mammal-eating) killer whales in the North Pacific also produce stereotyped whistles
82 and the repertoire seems to be shared by all members of the population (Riesch and
83 Deecke 2011). Despite being sympatric with resident killer whales, transients have
84 distinct, population-specific repertoires of both pulsed calls and whistles (Ford and
85 Fisher 1982, Riesch and Deecke 2011).

86 In Iceland and Norway, early studies suggested that killer whales also produce
87 group-specific call repertoires (Moore *et al.* 1988, Strager 1995). A common ecotype,
88 known as the “herring-eating Scandinavian killer whale” (Simon *et al.* 2007) has been
89 suggested for killer whales in this region due to close genetic relatedness (Foote *et al.*

2009b, Morin *et al.* 2010) and similar acoustic and foraging behavior. Killer whales are thought to have regularly migrated between these two locations (Jonsgård and Lyshoel 1970) due to the distribution of their main prey, Atlantic herring *Clupea harengus* (Sigurjónsson *et al.* 1988, Similä *et al.* 1996, Simon *et al.* 2007). The migration route of the Atlanto-Scandian herring stock spanned from western Norway to eastern Iceland prior to the stock collapse in the 1960's (Jakobsson and Østvedt 1999). The migration patterns of the Icelandic and Norwegian herring stocks changed following the collapse and remained closer to each country's coast (Jakobsson and Stefánsson 1999, Kvamme *et al.* 2003). Photo-identification studies dating to the 1980's show no matches between Iceland and Norway (Sigurjónsson *et al.* 1988, Foote *et al.* 2009a), suggesting that if killer whales migrated between those two locations in the past, there is currently little to no interaction between these populations. Nevertheless, some Icelandic killer whales have been re-identified as far east as Shetland (Foote *et al.* 2009a), showing that the movements of some individuals do extend beyond coastal waters. The suggested historic contact of killer whales between Iceland and Norway may have influenced their acoustic repertoire. However, comparisons of their pulsed call type repertoire have found apparently little repertoire sharing between the two locations (Moore *et al.* 1988, Strager 1995, Stenersen and Similä 2004, Shamir *et al.* 2014). If killer whales in Iceland and Norway were in contact in the past and their acoustic repertoires were similar, it is possible that sufficient time has passed for call repertoires to have diverged - explaining the pattern observed today. We might then expect that other signals produced by these whales may also show divergence. A divergent call repertoire may serve as a population-identifier, however, the extent of stereotypy and geographic variation in other signals is little understood.

High-frequency whistles ranging up to 75 kHz were first reported from herring-eating Northeast Atlantic killer whales (Samarra *et al.* 2010). High-frequency whistles have since been described from killer whale recordings made in the North Pacific (referred to as ‘high-frequency modulated signals’ and ‘ultrasonic whistles’; Simonis *et al.* 2012, Filatova *et al.* 2012) and in the Antarctic (Trickey *et al.* 2014), indicating that this is a widespread sound type produced by several populations although they do not appear to be ubiquitous (Samarra *et al.* 2010, Filatova *et al.* 2012). In the North Pacific, whistles were consistently downsweeps and, in some cases, highly stereotyped (Simonis *et al.* 2012, Filatova *et al.* 2012). Here, we analyze qualitatively and quantitatively the variation in time-frequency contours of high-frequency whistles recorded from herring-eating killer whales in Norway and Iceland, and compare them to stereotyped high-frequency whistles recorded in the North Pacific. Our objectives were: 1) to investigate the degree of micro and macro-geographic variation of time-frequency parameters and; 2) to investigate whether high-frequency whistles produced by killer whales in Iceland and Norway were also stereotyped and, if so, evaluate the spatial distribution of different whistle types. This study of geographic variation is a starting point to considering the potential function(s) of these poorly understood signals.

Methods

In the Northeast Atlantic, acoustic recordings were collected in Tysfjord, Vestfjord and Ofotfjord (Northern Norway) between 2005 and 2009 and off the Vestmannaeyjar archipelago and the Reykjanes peninsula in 2004, 2008 and 2009 (Iceland, Fig. 1). In both locations, recordings were collected during the day and most sounds were recorded from feeding or socializing whales. No other cetaceans were concurrently

sighted. Recording systems used varied with year and location and included towed and vertical hydrophone arrays, as well as sound recording tags attached to whales using suction cups (“Dtags”; Johnson and Tyack 2003). All systems sampled at 96 or 192 kHz (Table 1). Further details of all systems used and data collected are given in Samarra *et al.* (2010). High-frequency whistles analyzed here are the same as those reported in that study, including whistles with frequency contours at least partially below 48 kHz (up to 48 kHz whistles) and whistles with frequency contours entirely above 48 kHz (>48 kHz whistles), when available. Only 2.6% of whistles at least partially below 48 kHz recorded at 192 kHz sampling rate crossed 48 kHz, i.e. had minimum frequency below 48 kHz and maximum frequency above 48 kHz. This suggests that the lower sampling rate of 96 kHz likely resulted in a negligible loss of whistles with frequency contours at least partially below 48 kHz that could be sampled. High-frequency whistles were defined as signals with contours entirely above 17 kHz (Samarra *et al.*, 2010), as this was the maximum frequency previously reported for killer whale whistles (Thomsen *et al.* 2001).

To compare whistles produced in the Northeast Atlantic with those produced by killer whales in the North Pacific, we analyzed high-frequency whistles described by Simonis *et al.* (2012). The recordings were collected at sampling rates of 192 or 200 kHz from either a ship-based hydrophone array or a high-frequency acoustic recording package (HARP) and were scanned for high-frequency whistles through the use of Long-Term Spectral Averages (LTSAs) (Wiggins and Hildebrand 2007). To date, recordings from 19.9 cumulative years at 18 different sites across the North Pacific have been examined by analysts trained to recognize the presence of these signals (Fig. 1). There have been 19 acoustic encounters of killer whales at 9 different locations that included HFM signals, of which the temporal and spectral

characteristics of a subset from 6 locations are described here. The recording locations of signals described here include: Aleutian Islands, Washington Coast, Southern California Bight, Hoke Seamount, Kauai, and Pearl and Hermes Atoll (Table 1, Fig. 1). High-frequency whistles were attributed to killer whales by their clear association with other sounds known to be produced by killer whales or by visual observations of killer whales present in the area (Simonis *et al.* 2012). To ensure the data sets from the two ocean basins were comparable we only used signals detected in the North Pacific where contours were also entirely above 17 kHz. Table 1 shows the total number of whistles analyzed from each location. We pooled all data from the North Pacific for subsequent analyses, as sample sizes were too small to compare each location separately. Following inspection of recordings using Adobe Audition 2.0© (Blackmann-Harris window; FFT=2048 and 4096, for 96 kHz and 192 or 200 kHz sampling rates, respectively; 100% window width) whistle contours were traced from visual inspection of the spectrogram using a peaks contour extraction algorithm as developed by Buck and Tyack (1993; Hann window; frequency resolution=46.875 Hz; time resolution=0.667 ms). The following descriptive parameters were measured from the extracted fundamental frequency contour: start, half-way duration point (mid) and end frequency, minimum and maximum frequency, frequency range (maximum-minimum frequency) and duration.

All high-frequency whistles were assigned a quality score based on visual assessment of signal to noise ratio and overlap with other sounds, between 1 (poor, when the signal was barely detectable in the spectrogram) and 3 (high, when the full contour was clearly visible). Only clearly visible contours (of quality 3) with sufficient signal-to-noise ratio to extract the measurements required were extracted in our data sets from both ocean basins. It is likely that some whistles in our sample

were produced by the same group or the same individuals (mean \pm stdv [min; max] of number of whistles per recording day: 22.4 ± 22 [1; 95]). Photographs of whales were opportunistically collected while recordings were made in Iceland and Norway. Estimates of group size were not consistently collected across recording sessions in these two locations, but in general varied between 4-15 animals to large aggregations containing 50-100 individuals. Photo-identification was particularly challenging in Norway due to low-light conditions in winter and analysis has not been completed. In Iceland, on average 17 ± 14 [2; 41] individuals were identified per recording session and for those individuals that were resighted at least once (52 of 86) the number of resightings was 4 ± 2 [1; 9]. Recordings in the North Pacific were mostly from a single day in each location, and group sizes were generally not available (Table 1). It was not possible to identify which individuals produced high-frequency whistles recorded in this study, however, it is unlikely that the sampling procedure across different days or across wide geographic areas (such as for the North Pacific data) biased the repeated sampling of a small number of individuals within each location.

Stereotypy of high-frequency whistles

To investigate whether, like pulsed calls, high-frequency whistles fell into stereotyped categories, the data set was categorized using two different methods: 1) visual categorization by human observers as commonly used to identify whistle types and; 2) automated categorization using ARTwarp (Deecke and Janik 2006).

1) Categorization by a human observer

We first conducted visual classification by a human observer using the entire sample of high-frequency whistles to identify differences in the repertoires of different

whistle types across locations. Spectrograms of all whistles were generated in MATLAB (version 7.0.4; spectrogram parameters: FFT=2048 or 4096, for 96 kHz and 192 or 200 kHz sampling rates, respectively; overlap=87.5%; window function=Hann). The observer was blind to the origin of high-frequency whistles being classified. High-frequency whistles seemed to show continuous rather than discrete variation, and as a result were grouped into broad categories (e.g., Azevedo and Van Sluys 2005): upsweeps (rise in frequency with no inflection points), downsweeps (decrease in frequency with no inflection points), ascending-descending (rise in frequency followed by one inflection point and then a decrease in frequency), descending-ascending (decrease in frequency followed by one inflection point and then an increase in frequency), constant (small or no change in frequency and no inflection point), and other (if a contour did not fit any of the previous categories, Fig. 2).

2) Categorization using an automated method

To inspect variation at a finer scale, we conducted an independent automated categorization in MATLAB (version 7.11). A subset of the extracted fundamental frequency contours were analyzed by an adaptive resonance theory neural network that uses dynamic time-warping to calculate contour similarity (ARTwarp; Deecke and Janik 2006). Computation time constraints meant that it was impossible to input all extracted contours unless whistles were downsampled, reducing their time resolution. However, downsampling by a factor of 3 resulted in severe misclassification of the data (18 out of 19 categories included misclassified whistles) based upon visual inspection of generated categories. Therefore, the original time

239 resolution was used (0.667 ms), but only a subset of randomly chosen whistle
240 contours was categorized.

241 ARTwarp compares the similarity between an input contour and a contour that
242 defines each category (reference contour) to a user-defined similarity value, called the
243 vigilance parameter, to decide how each input contour should be categorized (Deecke
244 and Janik 2006). To find the categorization that would explain most variation in high-
245 frequency whistles with the least number of categories, the vigilance parameter was
246 set to values between 0% and 100% in 50 logarithmic steps, and the subset of
247 contours was categorized for each vigilance parameter value. The optimal
248 categorization should have the highest variance ratio, calculated as the ratio of
249 average within-category similarity over average between-category similarity as in
250 Deecke and Janik (2006). This method was adopted because it has successfully
251 classified the signature whistles of bottlenose dolphins and pulsed calls of killer
252 whales (Deecke and Janik 2006).

253

254 *Multivariate analysis of variation in time-frequency parameters*

255 We conducted a multivariate discriminant function analysis (DFA) to investigate
256 differences between whistles from different locations. All frequency and time
257 parameters were included, except frequency range, as it is already represented by
258 minimum and maximum frequency. We used location as the grouping variable
259 ('Iceland', 'Norway' or 'North Pacific') and used a jackknife cross-validation
260 technique implemented in the `lda` function of package MASS version 7.3-16
261 (Venables and Ripley 2002) in R 2.11.1 for Mac OS (R Development Core Team
262 2011). The overall proportion of correct classifications and the proportion of correct
263 classifications by location were calculated. These were compared to the proportion of

by-chance accuracy, calculated as the sum of the squares of all prior probabilities, assumed to be equal for all locations.

Results

We measured the parameters of 985 high-frequency whistles ($n_{\text{Iceland}} = 570$; $n_{\text{Norway}} = 256$; $n_{\text{Pacific}} = 159$). One of the whistles in our sample (from Norway) was a clear outlier, with duration of 4.2 s, much longer than the majority of remaining whistles and was removed from the analyses. Table 2 presents the descriptive statistics of all contour parameters measured. Duration and frequency range showed the highest coefficients of variation across all locations. All other frequency parameters had much lower coefficients of variation, however variation for Iceland and Norway was consistently larger than variation in the North Pacific. This could be due to the smaller sample size of whistles from the Pacific. However, in the North Atlantic very high-frequency whistles were recorded that likely explain the larger coefficients of variation observed here. There was a clear gap in the distribution of frequency parameters between whistles with fundamental contours up to 48 kHz and whistles with fundamental frequency contours entirely above 48 kHz (Fig. 3). Whistles with contours entirely above 48 kHz were recorded in a small number of encounters ($n_{\text{Iceland}} = 4$; $n_{\text{Norway}} = 2$). They appeared as clear outliers in the distributions and were thus removed from the comparisons of parameter distributions (Fig. 3).

There was considerable overlap between the distributions of some of the contour parameters from Iceland and Norway (Fig. 3). To compare these parameters across locations, we employed a Mann-Whitney U test, due to the non-normality of most distributions (Shapiro-Wilk normality tests: $P < 0.0001$, except for frequency range in

the North Pacific with $p=0.09$). We used Bonferroni correction to adjust the significance level to account for multiple comparisons ($0.05/21=0.002$). Despite the apparent overlap, comparisons between the parameter distributions from Iceland and Norway revealed significant differences in end frequency (Mann-Whitney U test: $W=74148$; $P=0.0004$), frequency range (Mann-Whitney U test: $W=81867$; $P<0.0001$) and maximum frequency (Mann-Whitney U test: $W=74203$; $P=0.0003$). End frequency and maximum frequency were significantly correlated within the two locations (Pearson correlation; Iceland: end vs. maximum frequency: $r=0.96$, $P<0.001$; Norway: end vs. maximum frequency: $r=0.98$, $P<0.001$), which likely reflects a whistle upsweep shape. Whistles recorded in the North Pacific tended to have lower start, mid, end, minimum, and maximum frequencies, a slightly longer duration and a greater frequency range. The start frequency was not significantly different between Iceland and the Pacific (Mann-Whitney U test: $W=49578$; $P=0.008$), while all other parameters were (P -values <0.0001). All parameters were also significantly different between the North Pacific and Norway, with the exception of duration (Mann-Whitney U test: $W=16777$; $P=0.11$).

305

306 *Categorization by a human observer*

The visual classification of whistles showed that in Iceland the majority of whistles were upsweeps, followed by descending-ascending whistle types, while in Norway, upsweeps and descending-ascending whistles were equally common (Table 3). Overall, the different whistle types recorded from Norway and Iceland largely resembled each other in spectral and temporal characteristics (Fig. 2), in agreement with the similarities suggested by the overlapping parameter distributions. In contrast, whistles from the North Pacific were all downsweeps, with only one whistle being

classified as ‘Other’. These clear differences in repertoire correspond to the differences in extracted frequency parameters (above). Nevertheless, the simple structure of downsweeps recorded in the North Pacific was similar to that of the few downsweeps recorded in the Northeast Atlantic (Fig. 2).

Categorization using an automated method

An assessment of the geographic variation of whistles was also completed using 50 randomly chosen whistles from each location. The results of the ARTwarp categorization showed that an increase in the vigilance parameter resulted in an increase in the number of categories generated. The chosen categorization divided the 150 high-frequency whistles into 27 categories, containing between 1 and 18 contours (mean \pm standard deviation of contours in each category: 6 ± 4.6). This categorization corresponded to the local maximum in variance ratio achieved when the vigilance parameter was 95.6%. All but five categories included more than one whistle. Of the five categories containing only one whistle one category contained one whistle from the Pacific, two categories contained single whistles from Norway, and two categories contained single whistles from Iceland. Of the 22 categories with more than one whistle, five included only high-frequency whistles from the Pacific. The remaining high-frequency whistles from the Pacific were grouped into three more categories that also included whistles from Norway: two categories containing only two whistles (one from the Pacific and one from Norway) and; one category including 17 whistles from the Pacific and one whistle from Norway. All remaining categories included whistles from both Norway and Iceland.

338 *Multivariate analysis of variation in time-frequency parameters*

339 The multivariate DFA showed high macrogeographic variation between the
340 Northeast Atlantic and North Pacific groups, with the first discriminant function
341 accounting for 98.9% of between group variability. The loadings of the first
342 discriminant function revealed that duration, maximum frequency, and end frequency
343 were the main discriminating predictors (Loadings: duration = -2.81; maximum
344 frequency = 0.33; end frequency = -0.46). The cross-validated classification showed
345 an overall correct classification of 62.4% of whistles to the correct location, compared
346 to a by-chance proportion of 33.3%, calculated as the sum of the squares of all prior
347 probabilities. Correct classification scores per location revealed that the North Pacific
348 was the location with the highest score (95.0%), with only 8 of its 159 whistles
349 misclassified as being from Norway. Lower correct classification scores (63.9% and
350 52.4%) were achieved for whistles from Norway and Iceland, respectively. Most
351 misclassifications for Norway (77 of 84) were assigned to Iceland and conversely,
352 most misclassifications for Iceland (254 of 261) were assigned to Norway. The
353 remaining 7 misclassifications from each location were assigned to the North Pacific.
354 Figure 4 shows the first two discriminant functions and illustrates how the first
355 discriminant function achieved the most discrimination between locations, with
356 whistles from the North Pacific being clearly different from those recorded in the
357 Northeast Atlantic, while whistles from Norway and Iceland overlapped to a much
358 greater extent. However, at high values of the second discriminant function there was
359 little overlap between Iceland and Norway (Fig. 4). Inspection of these whistles
360 revealed that these have high frequency range and possibly are driving observed
361 statistically significant differences in this parameter between locations.

To investigate whether discrimination between Iceland and Norway was more apparent within whistle types, discriminant function analyses were carried out for the two most common whistle types (upsweeps and descending-ascending) at these locations. Descending-ascending whistles were correctly classified at rates of 60% and 66.7% for Iceland and Norway, respectively. Overall correct classification was of 62.7%, compared to a by-chance proportion of 50%, suggesting a slight improvement in classification when using this whistle type, but still considerable similarity between locations. The overlap in the distribution of the discriminant scores resulting from the first discriminant function is presented in Figure 5, illustrating how the discrimination between these two locations was poor. Correct classification of upsweeps was 58.7% and 58.5% for Iceland and Norway, respectively. For upsweeps, overall correct classification was 58.6%, compared to a by-chance proportion of 50%, which also suggests an overlap in the discriminant scores within this type (Fig. 5).

Discussion

We found clear macrogeographic variation between high-frequency whistles recorded in the North Pacific and the Northeast Atlantic, while within the Northeast Atlantic only subtle microgeographic variation was observed. In all locations whistles could be manually classified into broad stereotyped categories, but the diversity of whistle types identified varied between ocean basins. While the majority of whistles recorded in the North Pacific consisted of only one whistle type, the Northeast Atlantic repertoires included more types and the repertoires were similar between locations. In all locations the within-location variation in most frequency parameters was small in comparison to variation in frequency range and duration. It is possible that some degree of this variation is due to varied signal-to-noise ratio conditions

under which recordings were collected. Nevertheless, in other delphinids intra-specific variation in frequency parameters was generally low, while variation in whistle duration tended to be comparatively higher (e.g., Ding *et al.* 1995b, Morisaka *et al.* 2005). It is possible that frequency parameters are constrained by morphology, while the variability of whistle duration may be related to social and non-social behavior or environmental factors, or random individual variation (e.g., May-Collado and Wartzok 2008).

Macrogeographic variation in high-frequency whistles

Within the North Pacific, acoustic encounters of killer whales were not common at any of the recording sites, and high-frequency whistles were infrequently observed. There was broad similarity in the recorded whistles despite the considerable geographic range covered by the recordings. The consistency in whistle type (downsweeps) and similarities in frequency characteristics to those reported by Filatova *et al.* (2012) suggests that within the Pacific Ocean there may be little variability in high-frequency whistles. However, not all populations appear to produce these signals; both Northeast Pacific resident and transient killer whales apparently do not produce high-frequency whistles (Samarra *et al.* 2010, Filatova *et al.* 2012) but the ecotypes of those whales that do produce them in the North Pacific remain largely unknown, with the exception of North Pacific offshores (Simonis *et al.* 2012, Filatova *et al.* 2012). While Pacific offshore, resident and Northeast Atlantic killer whales are all closely related genetically (Morin *et al.* 2010), it is curious that Pacific resident killer whales have not been recorded using the signals shared by their nearest relatives. In contrast to the apparent similarity within the Pacific Ocean, there were clear differences between whistles recorded in the Pacific and Atlantic Ocean basins.

412 Whistles recorded in the North Pacific had consistently lower frequency
413 parameters, while frequency range was significantly higher in the North Pacific than
414 in either of the Northeast Atlantic populations. Differences in duration were less
415 pronounced and significant differences were only detected in comparison to Iceland.
416 The discriminant function analysis was able to correctly classify the vast majority of
417 whistles from the North Pacific, assigning duration and maximum and end frequency
418 as main discriminating predictors. Whistle types identified were also considerably
419 different between ocean basins; while downsweeps were the most common whistle
420 type in the North Pacific, this whistle type was uncommon in the Northeast Atlantic.
421 The automated categorization also grouped most whistles from the Pacific into
422 distinct categories. This divergence in whistle types will likely influence some of the
423 observed differences in frequency parameters particularly the start, mid and end
424 frequency but does not explain differences in minimum and maximum frequencies,
425 which should not be affected by whistle shape. Thus, we believe that the observed
426 differences in frequency parameters between ocean basins are not exclusively due to
427 differences in whistle type usage but reflect a real divergence in the whistle frequency
428 produced.

429 Divergence in frequency at macrogeographic scales could reflect divergence in
430 geographically isolated populations that could originate from a wide range of genetic
431 and social mechanisms. Indeed genetic data suggests considerable variation between
432 populations in the Atlantic and the Pacific in comparison to variation between Iceland
433 and Norway (Morin *et al.* 2010, Foote *et al.*, 2011). Differences in the acoustic
434 environment, such as background noise or transmission properties (*e.g.*, Morisaka *et*
435 *al.* 2005, May-Collado and Wartzok 2008), have also been proposed as factors
436 explaining variations in signal frequency characteristics between populations. In both

ocean basins recordings were collected over wide geographic areas, and we therefore cannot identify large and consistent habitat differences that could clearly explain the patterns of variation observed in this study. Body size, however, is known to vary between the two ocean basins; killer whales in the Northeast Atlantic in general have smaller body sizes than killer whales in the Northeast Pacific (Christensen 1984, Stenersen and Similä 2006), but Northeast Pacific offshores have smaller body size than resident and transient killer whales (Ford *et al.* 2000, Dahlheim *et al.* 2008). Nevertheless, we cannot exclude the possibility that divergence in frequency characteristics between these ocean basins may be related to morphological constraints. The reasons behind the consistent production of downsweeps in the North Pacific in contrast to the more variable repertoires recorded in the Northeast Atlantic are unknown. Although intriguing, to fully understand the reasons behind variations in frequency and usage of whistle types across ocean basins more effort is required to record killer whales with adequate sampling rates in other locations. This will reveal the extent of the consistent downsweep repertoire across the North Pacific or the existence of variable repertoires in other ocean basins as well as how frequency characteristics may vary in other habitats.

Microgeographic variation in high-frequency whistles

Within the Northeast Atlantic, we found similarities in the repertoire and characteristics of high-frequency whistles produced by killer whales in Norway and Iceland. None of the quantitative methods employed was able to distinguish between whistles from Iceland and Norway as clearly as between whistles from Northeast Atlantic and North Pacific, suggesting different levels of divergence between locations. Despite the apparent overlap in parameter distributions in the Northeast

462 Atlantic, whistles from Norway showed significantly lower end frequency, maximum
463 frequency and frequency range when compared to Iceland. The discriminant function
464 analysis showed some degree of correct classifications, although many whistles from
465 both locations were misclassified, while the automated categorization included
466 whistles from both locations in most of the categories generated. Although the overall
467 repertoire was similar between the two locations, there were some differences in the
468 usage of whistle types. Upsweeps were the most common whistle type in Iceland,
469 while in Norway both upsweeps and descending-ascending whistles were equally
470 common. Even when trying to discriminate between locations within whistles of the
471 same type, discriminant function analyses still misclassified a large proportion of
472 whistles, suggesting similarity between whistles produced. The similarity in both
473 time-frequency parameters and overall repertoire of high-frequency whistles between
474 Norway and Iceland is in striking contrast to the divergence in pulsed call repertoires
475 (Moore *et al.* 1988, Strager 1995, Stenersen and Similä 2004, Shamir *et al.* 2014). At
476 present we have little evidence of contextual production of high-frequency whistles to
477 help us identify their function. Nevertheless, this divergence between pulsed calls and
478 high-frequency whistles possibly reflects different functions of these different signal
479 types.

480 Microgeographic variation in acoustic signals may be shaped by genealogy, the
481 timing of separation, ranging behavior of individuals, or cultural divergence in the
482 case of learned acoustic signals (*e.g.*, Ding *et al.* 1995b, Azevedo and Van Sluys
483 2005, Papale *et al.* 2013). For killer whales in Norway and Iceland evidence for (a
484 recent) common ancestor comes from historic ranging patterns of prey, genetics
485 (Jonsgård and Lyshoel 1970, Foote *et al.* 2009a), and high-frequency whistle
486 repertoires, while the unique pulsed call repertoires of each region may reflect more

487 recent divergence (Moore *et al.* 1988, Strager 1995, Stenersen and Similä 2004,
488 Shamir *et al.* 2014). A small number of high-frequency whistles have also been
489 recorded from killer whales in Shetland (Samarra *et al.* 2010), an archipelago situated
490 between Iceland and Norway where some Icelandic killer whales are known to travel
491 (Foote *et al.* 2009a). Further recordings from this location and others should provide
492 an interesting comparison to evaluate the degree of divergence between adjacent
493 populations. Although the function(s) of these high-frequency whistles remain
494 unclear, the signals analyzed in this study offer a window to understanding how
495 acoustic behavior may relate to ancestry and dispersal patterns of killer whale
496 populations on multiple scales.

497

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522

523 **Literature cited**

- 524 Azevedo, A. F. and M. Van Sluys. 2005. Whistles of tucuxi dolphins (*Sotalia*
525 *fluviatilis*) in Brazil: comparisons among populations. Journal of the Acoustical
526 Society of America 117:1456-1464.
- 527 Baron, S. C., A. Martinez, L. P. Garrison and E. O. Keith. 2008. Differences in
528 acoustic signals from delphinids in the western North Atlantic and northern Gulf
529 of Mexico. Marine Mammal Science, 24(1): 42-56.
- 530 Buck, J. R. and P. L. Tyack. 1993. A quantitative measure of similarity for *Tursiops*
531 *truncatus* signature whistles. Journal of the Acoustical Society of America
532 94:2497-2506.
- 533 Caldwell, M. C. and D. K. Caldwell. 1968. Vocalization of naïve dolphins in small
534 groups. Science 159:1121-1123.
- 535 Camargo, F. S., M. M. Rollo, V. Giampaoli and C. Bellini. 2006. Whistle variability
536 in South Atlantic spinner dolphins from the Fernando de Noronha Archipelago
537 off Brazil. Journal of the Acoustical Society of America 120(6):4071-4079.
- 538 Christensen, I. 1984. Growth and reproduction of killer whales, *Orcinus orca*, in
539 Norwegian coastal waters. Report of the International Whaling Commission
540 6:253-258.
- 541 Dahlheim, M. E., Schulman-Janiger, A., Black, N., Ternullo, R., Ellifrit, D. and K. C.
542 Balcomb. 2008. Eastern temperate North Pacific offshore killer whales (*Orcinus*
543 *orca*): occurrence, movements, and insights into feeding ecology. Marine
544 Mammal Science 24(3):719-729.
- 545 Deecke, V. B. and V. M. Janik. 2006. Automated categorization of bioacoustic
546 signals: Avoiding perceptual pitfalls. Journal of the Acoustical Society of
547 America 119:645-653.

- 548 Deecke, V.B., L. G. Barrett-Lennard, P. Spong and J. K. B. Ford. 2010. The structure
549 of stereotyped calls reflects kinship and social affiliation in resident killer whales
550 (*Orcinus orca*). *Naturwissenschaften* 97:513–518.
- 551 Ding, W., B. Würsig and W. Evans. 1995a. Comparisons of whistles among seven
552 odontocete species. Pages 299-323 in R. A. Kastelein, J. A. Thomas and P. E.
553 Nachtigall, eds. *Sensory systems of aquatic mammals*. De Spil Publishers,
554 Woerden, The Netherlands.
- 555 Ding, W., B. Würsig and W. E. Evans. 1995b. Whistles of bottlenose dolphins:
556 comparisons among populations. *Aquatic Mammals* 21.1:65-77.
- 557 Filatova, O.A., J. K. B. Ford, C. O. Matkin, L. G. Barrett-Lennard, A. M. Burdin and
558 E. Hoyt. 2012. Ultrasonic whistles of killer whales (*Orcinus orca*) recorded in the
559 North Pacific (L). *Journal of the Acoustical Society of America* 132:3618-3621.
- 560 Ford, J. K. B. and H. D. Fisher. 1982. Killer whale (*Orcinus orca*) dialects as an
561 indicator of stocks in British Columbia. *Reports of the International Whaling*
562 *Commission* 32:671-679.
- 563 Ford, J. K. B. 1989. Acoustic behavior of resident killer whales (*Orcinus orca*) off
564 Vancouver Island, British Columbia. *Canadian Journal of Zoology* 67, 727-745.
- 565 Ford, J. K. B. 1991. Vocal traditions among resident killer whales (*Orcinus orca*) in
566 coastal waters of British Columbia. *Canadian Journal of Zoology* 69:1454-1483.
- 567 Ford, J. K. B., Ellis, G. M. and K. C. Balcomb. 2000. *Killer whales: the natural*
568 *history and genealogy of Orcinus orca in British Columbia and Washington*.
569 Second Edition. Vancouver: University of British Columbia Press.
- 570 Foote, A. D., T. Similä, G. A. Vikingsson and P. T. Stevick. 2009a. Movement, site
571 fidelity and connectivity in a top marine predator, the killer whale. *Evolutionary*
572 *Ecology* 24:803-814.

- 573 Foote, A. D., J. Newton, S. B. Piernney, E. Willerslev and M. T. P. Gilbert. 2009b.
574 Ecological, morphological and genetic divergence of sympatric North Atlantic
575 killer whale populations. *Molecular Ecology* 18:5207-5217.
- 576 Foote, A. D., J. T. Vilstrup, R. de Stephanis, *et al.* 2011. Generic differentiation
577 among North Atlantic killer whale populations. *Molecular Ecology* 20(3):629-
578 641.
- 579 Jakobsson, J. and O. J. Østvedt. 1999. A review of joint investigations on the
580 distribution of herring in the Norwegian and Iceland Seas 1950-1970. *Rit*
581 *Fiskideildar* 16:209-238.
- 582 Jakobsson, J. and G. Stefánsson. 1999. Management of summer-spawning herring off
583 Iceland. *ICES Journal of Marine Science* 56:827-833.
- 584 Janik, V. M., L. S. Sayigh and R. S. Wells. 2006. Signature whistle shape conveys
585 identity information to bottlenose dolphins. *Proceedings of the National Academy*
586 *of Sciences* 103:8293-8297.
- 587 Janik, V. M. 2009. Acoustic communication in delphinids. *Advances in the Study of*
588 *Behavior* 40:123-157.
- 589 Johnson, M. P. and P. L. Tyack. 2003. A digital acoustic recording tag for measuring
590 the response of wild marine mammals to sound. *IEEE Journal of Oceanic*
591 *Engineering* 28:3-12.
- 592 Jonsgård, Å. And P. B. Lyshoel. 1970. A contribution to the knowledge of the biology
593 of the killer whale *Orcinus orca* (L.). *Nytt Magasin for Zoologi* 18:41-48.
- 594 Kvamme, C., L. Nøttestad, A. Fernö, *et al.* 2003. Migration patterns in Norwegian
595 spring-spawning herring: why young fish swim away from the wintering area in
596 late summer. *Marine Ecology Progress Series* 247:197-210.

- 597 May-Collado, L. J. and D. Wartzok. 2008. A comparison of bottlenose dolphin
598 whistles in the Atlantic Ocean: factors promoting whistle variation. *Journal of*
599 *Mammalogy* 89(5):1229-1240.
- 600 Matthews, J. N., L. E. Rendell, J. C. D. Gordon and D. W. Macdonald. 1999. A
601 review of frequency and time parameters of cetacean tonal calls. *Bioacoustics*
602 10:47-71.
- 603 Miller, P. J. O. and D. E. Bain. 2000. Within-pod variation in the sound production of
604 a pod of killer whales, *Orcinus orca*. *Animal Behaviour* 60:617-628.
- 605 Moore, S. E., J. K. Francine, A. E. Bowles and J. K. B. Ford. 1988. Analysis of calls
606 of killer whales, *Orcinus orca*, from Iceland and Norway. *Rit Fiskideildar*
607 11:225-250.
- 608 Morin, P. A., F. I. Archer, A. D. Foote, *et al.* 2010. Complete mitochondrial genome
609 phylogeographic analysis of killer whales (*Orcinus orca*) indicates multiple
610 species. *Genome Research* 20:908-916.
- 611 Morisaka, T., M. Shinohara, F. Nakahara and T. Akamatsu. 2005. Geographic
612 variations in the whistles among three Indo-Pacific bottlenose dolphin *Tursiops*
613 *aduncus* populations in Japan. *Fisheries Science* 71:568-576.
- 614 Munding, P. C. 1982. Microgeographic and macrogeographic variation in the
615 acquired vocalizations of birds. Pages 147-208 *in* D. E. Kroodsma, E. H. Miller,
616 and H. Ouellet, eds. *Acoustic Communication in Birds: Song learning and its*
617 *consequences*. Vol. 2, Academic Press, New York, NY.
- 618 Nousek, A. E., P. J. B. Slater, C. Wang and P. J. O. Miller. 2006. The influence of
619 social affiliation on individual vocal signatures of northern resident killer whales
620 (*Orcinus orca*). *Biology Letters* 2:481-484.

- 621 Papale, E., M. Azzolin, I. Cascão, *et al.* (2014). Macro- and micro-geographic
622 variation of short-beaked common dolphin's whistles in the Mediterranean Sea
623 and Atlantic Ocean. *Ethology, Ecology and Evolution* 26:392-404.
- 624 Podos, J., V. M. F. da Silva and M. R. Rossi-Santos. 2002. Vocalizations of Amazon
625 river dolphins, *Inia geoffrensis*: insights into the evolutionary origins of delphinid
626 whistles. *Ethology* 108:601-612.
- 627 R Development Core Team. 2011. R: A language and environment for statistical
628 computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-
629 900051-07-0, URL <http://www.R-project.org/>.
- 630 Rendell, L E., J. N. Matthews, A. Gill, J. C. D. Gordon and D. W. Macdonald. 1999.
631 Quantitative analysis of tonal calls from five odontocete species, examining
632 interspecific and intraspecific variation. *Journal of Zoology* 249:403-410.
- 633 Riesch, R., J. K. B. Ford and F. Thomsen. 2006. Stability and group specificity of
634 stereotyped whistles in resident killer whales, *Orcinus orca*, off British Columbia.
635 *Animal Behaviour* 71:79-91.
- 636 Riesch, R., J. K. B. Ford and F. Thomsen. 2008. Whistle sequences in wild killer
637 whales (*Orcinus orca*). *Journal of the Acoustical Society of America* 124:1822-
638 1829.
- 639 Riesch, R. and V. B. Deecke. 2011. Whistle communication in mammal-eating killer
640 whales (*Orcinus orca*): further evidence for acoustic divergence between
641 ecotypes. *Behavioral Ecology and Sociobiology* 65:1377-1387.
- 642 Rossi-Santos, M. R. and J. Podos. 2006. Latitudinal variation in whistle structure of
643 the estuarine dolphin *Sotalia guianensis*. *Behaviour* 143:347-364.

- 644 Samarra, F. I. P., V. B. Deecke, K. Vinding, M. H. Rasmussen, R. J. Swift and P. J.
645 O. Miller. 2010: Killer whales (*Orcinus orca*) produce ultrasonic whistles. Journal
646 of the Acoustical Society of America 128:EL205-EL210.
- 647 Sigurjónsson, J., T. Lyrholm, S. Leatherwood, E. Jónsson and G. Víkingsson. 1988.
648 Photoidentification of killer whales, *Orcinus orca*, off Iceland, 1981 through
649 1986. Rit Fiskideildar 11:99-114.
- 650 Similä, T., J. C. Holst and I. Christensen. 1996. Occurrence and diet of killer whales
651 in northern Norway: seasonal patterns relative to the distribution and abundance
652 of Norwegian spring-spawning herring. Canadian Journal of Fisheries and
653 Aquatic Sciences 53:769-779.
- 654 Simon, M., P. K. McGregor and F. Ugarte. 2007. The relationship between the
655 acoustic behaviour and surface activity of killer whales (*Orcinus orca*) that feed
656 on herring (*Clupea harengus*). Acta Ethologica 10:47-53.
- 657 Simonis, A. E., S. Baumann-Pickering, E. Oleson, M. L. Melcón, M. Gassmann, S.
658 M. Wiggins and J. A. Hildebrand. 2012. High-frequency modulated signals of
659 killer whales (*Orcinus orca*) in the North Pacific. Journal of the Acoustical
660 Society of America 131:EL295-EL301.
- 661 Shamir, L., C. Yerby, R. Simpson, A. M. von Benda-Beckmann, P. Tyack, F.
662 Samarra, P. Miller and J. Wallin. 2014. Classification of large acoustic datasets
663 using machine learning and crowdsourcing: application to whale calls. Journal of
664 the Acoustical Society of America, 135(2):953-962.
- 665 Steiner, W. W. 1981. Species-specific differences in pure tonal whistle vocalizations
666 of five Western North Atlantic dolphin species. Behavioral Ecology and
667 Sociobiology 9(4):241-246.

- 668 Stenersen, J. and T. Similä. 2004. Norwegian killer whales. Henningsvær: Tringa
669 Forlag.
- 670 Strager, H. 1995. Pod-specific call repertoires and compound calls of killer whales,
671 *Orcinus orca*, Linnaeus, 1758, in the waters of northern Norway. Canadian
672 Journal of Zoology 73:1037-1047.
- 673 Thomsen, F., D. Franck and J. K. B. Ford. 2001. Characteristics of whistles from the
674 acoustic repertoire of resident killer whales (*Orcinus orca*) off Vancouver Island,
675 British Columbia. Journal of the Acoustical Society of America 109:1240-1246.
- 676 Thomsen, F., D. Franck and J. K. B. Ford. 2002. On the communicative significance
677 of whistles in wild killer whales (*Orcinus orca*). Naturwissenschaften 89:404-407.
- 678 Trickey, J. S., M. V. R. Reyes, S. Baumann-Pickering, M. L. Melcón, J. A.
679 Hildebrand and M. A. Iñíguez. 2014. Acoustic encounters of killer and beaked
680 whales during the 2014 SORP cruise. IWC report SC/65b/SM12.
- 681 Tyack, P. 1986. Population Biology, Social behavior and communication in whales
682 and dolphins. Trends in Ecology and Evolution 1:144-150.
- 683 Van Parijs, S. M. and P. J. Corkeron. 2001. Vocalizations and behaviour of Pacific
684 humpback dolphins *Sousa chinensis*. Ethology 107:701-716.
- 685 Venables, W. N. and B. D. Ripley. 2002. Modern applied statistics with S. Fourth
686 Edition. Springer, New York. NY.
- 687 Wiggins, S. M. and J. A. Hildebrand. 2007. High-frequency acoustic recording
688 package (HARP) for broad-band, long-term marine mammal monitoring. Pages
689 551-557 in International Symposium on Underwater Technology and Workshop
690 on Scientific Use of Submarine Cables and Related Technologies (2007).

691 **Table 1.** Specifications of recordings containing high-frequency whistles analyzed in this study (further details are given in Samarra *et al.* 2010
 692 and Simonis *et al.* 2012).

Ocean basin	Location	Recording days	Recording time (h)	Sampling rate (kHz)	Recording method	Whistles analyzed
Northeast Atlantic	Iceland	19	64	96 and 192	Vertical hydrophone array	570
					Towed hydrophone array	
					Dtag	
North Pacific	Norway	18	104	96 and 192	Towed hydrophone array	257
					Dtag	
	Aleutian islands	2	3.1	200	HARP (depth 783 m)	44
	Hoke Seamount	1	0.6	200	HARP (depth 770 m)	22
	Kauai	1	0.6	200	HARP (depth 706 m)	36
	Pearl and Hermes Atoll	1	0.6	200	HARP (depth 753 m)	10
	Southern California	1	1.9	200	HARP (depth 1295 m)	22

Bight						
Southern California	1	0.5	200	Ship-based hydrophone	11	
Bight				array		
Washington Coast	1	0.7	192	Ship-based hydrophone	14	
				array		

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694

Table 2. Descriptive statistics of high-frequency whistles analyzed in this study. Sample sizes for each location are included in brackets. For each parameter, values given are mean \pm standard deviation [minimum – maximum], with coefficient of variation (calculated as the ratio of the standard deviation to the mean) given as a percentage within brackets. All frequency parameters are presented in kHz and duration is presented in milliseconds (ms).

Location	Iceland (<i>n</i> =570)	Norway (<i>n</i> =256)	North Pacific (<i>n</i> = 159)
Start frequency	32.6 \pm 9.1 (27.9%) [16.9 – 71.2]	34.7 \pm 11.0 (31.7%) [18.3 – 71.0]	29.6 \pm 5.1 (17.2%) [19.3 – 44.0]
End frequency	38.2 \pm 8.7 (22.8%) [19.4 – 74.7]	37.4 \pm 9.0 (24.1%) [21.5 – 68.3]	20.8 \pm 3.2 (15.4%) [17.1 – 33.4]
Mid frequency	33.8 \pm 8.6 (25.4%) [17.6 – 68.8]	34.6 \pm 9.6 (27.7%) [19.0 - 64.3]	25.3 \pm 4.3 (16.9%) [18.4 – 39.4]
Minimum frequency	31.7 \pm 8.6 (27.1%) [16.9 - 68.3]	33.0 \pm 9.2 (27.9%) [18.1 – 64.3]	20.8 \pm 3.2 (15.3%) [17.1 – 33.4]
Maximum frequency	38.4 \pm 8.7 (22.7%) [19.4 – 74.7]	38.4 \pm 10.2 (26.6%) [22.3 – 71.0]	29.7 \pm 5.1 (17.1%) [19.3 – 44.0]
Frequency range	6.8 \pm 3.7 (54.4%)	5.4 \pm 2.9 (53.7%)	8.9 \pm 3.8 (42.6%)

	[0.7 – 21.2]	[1.0 – 19.9]	[1.6 – 20.2]
Duration	138.1 ± 135.9	143.6 ± 143.1	142.6 ± 74.2
	(98.4%)	(99.7%)	(52.1%)
	[6 - 814]	[10 - 1300]	[37.8 – 371.2]

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For Peer Review

703 **Table 3.** Proportion of high-frequency whistle types produced in different locations.

704 Desc-Asc stands for descending-ascending and Asc-Desc stands for ascending-

705 descending whistles.

Whistle categories							
Location	Upsweep	Downsweep	Desc-Asc	Asc-Desc	Constant	Other	Total
Iceland	68.4%	1.4%	28.6%	0%	0.2%	1.4%	570
Norway	41.2%	9.3%	42.8%	0.8%	0%	5.8%	257
North Pacific	0%	99.4%	0%	0%	0%	0.6%	159
Total	50.3%	19.3%	27.7%	0.2%	0.1%	2.4%	986

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Figure captions

Figure 1. Map illustrating the approximate locations (stars) where recordings containing high-frequency whistles were collected in: top) the Northeast Atlantic and; bottom) the North Pacific. SCB stands for Southern California Bight. Locations where recordings were conducted but no high-frequency whistles were detected are also shown (circles).

Figure 2. Spectrograms showing examples of different high-frequency whistle types from Iceland, Norway and the North Pacific. If a specific whistle type was only produced in one location only one example was shown. Note the different y-axis scaling for the Constant whistle type.

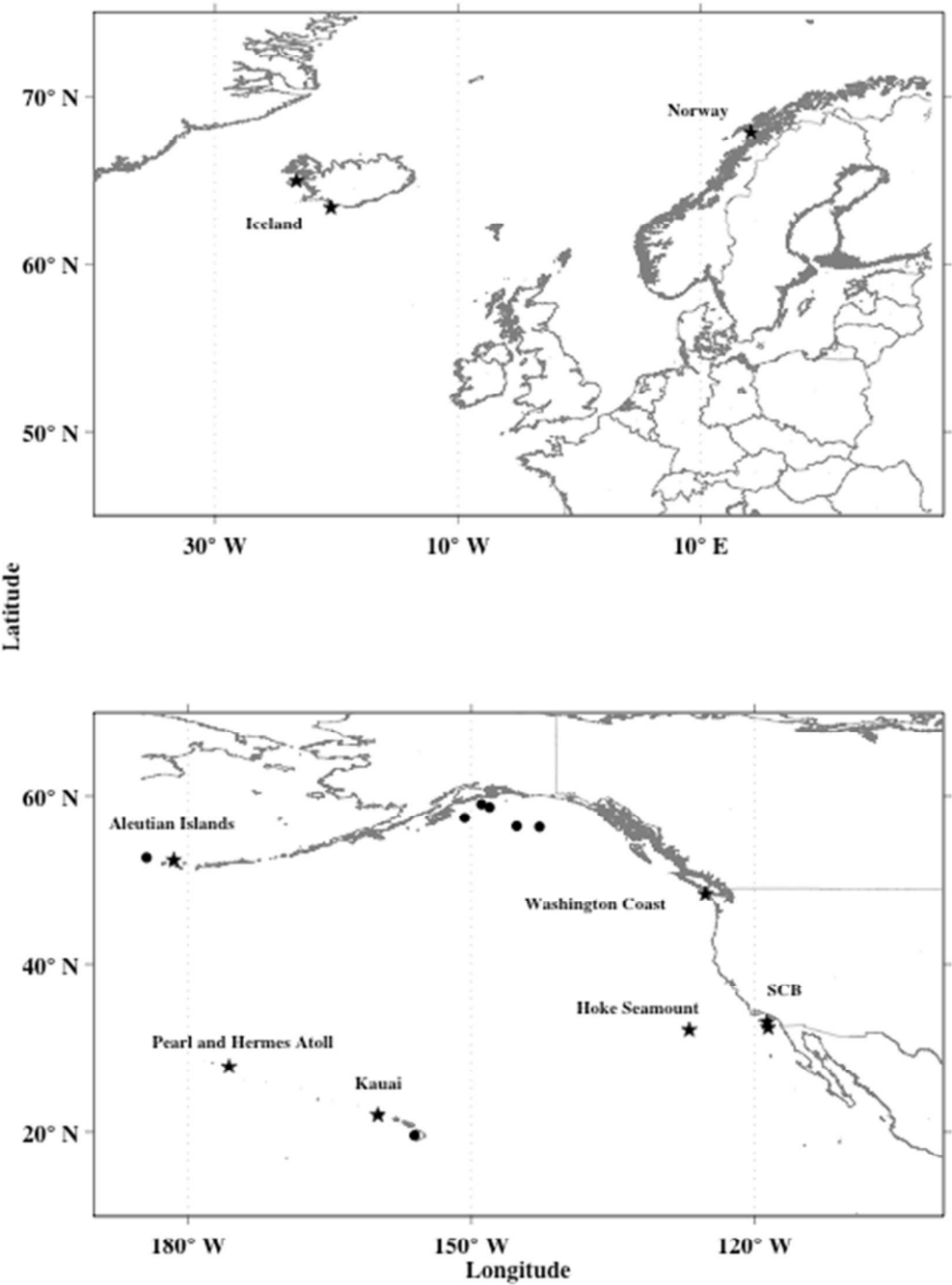
Figure 3. Distribution of all frequency parameters extracted from whistle contours. Horizontal lines represent medians, boxes represent inter-quartiles, and whiskers represent values within 1.5 times the inter-quartile range from the boxes. Outliers are plotted as single points. Frequency range is plotted separately due to its different y-axis scale, as is duration.

Figure 4. Plot of the first two discriminant functions for the comparison between whistles recorded in Iceland (I), Norway (N) and the North Pacific (P). Colors follow the same legend as in Figure 2. Note the overlap between whistles from Norway (N) and Iceland (I) in contrast to discrimination between whistles from the NE Atlantic and whistles from the Pacific (P).

732 **Figure 5.** Boxplots illustrating the distribution of discriminant scores (DF1) for
733 descending-ascending whistles (Desc-Asc) for the two groups (Iceland and Norway)
734 and discriminant scores for upsweep whistles for the two groups. Colors follow the
735 same legend as in Figure 2.
736

For Peer Review

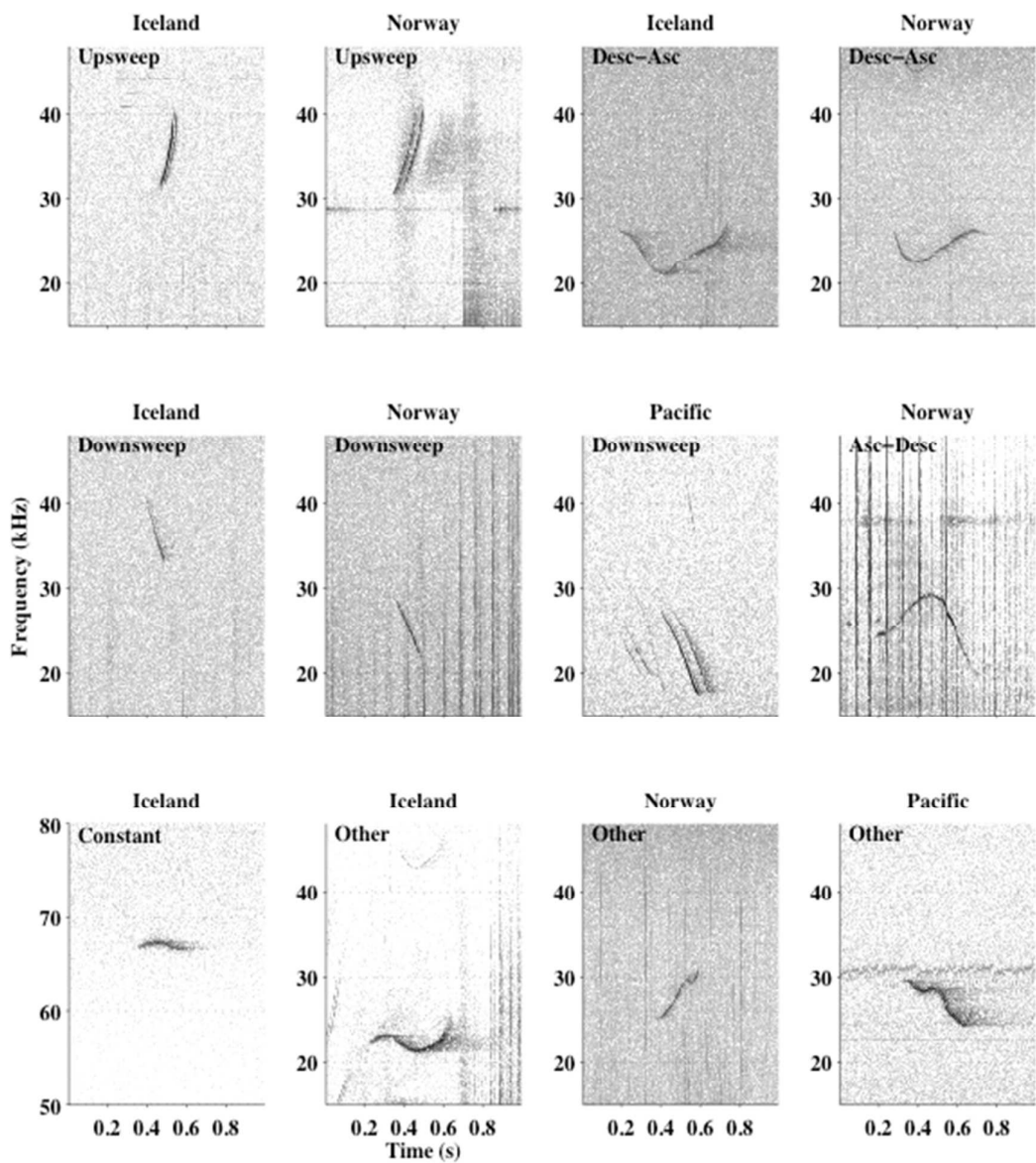
737 **Figure 1**



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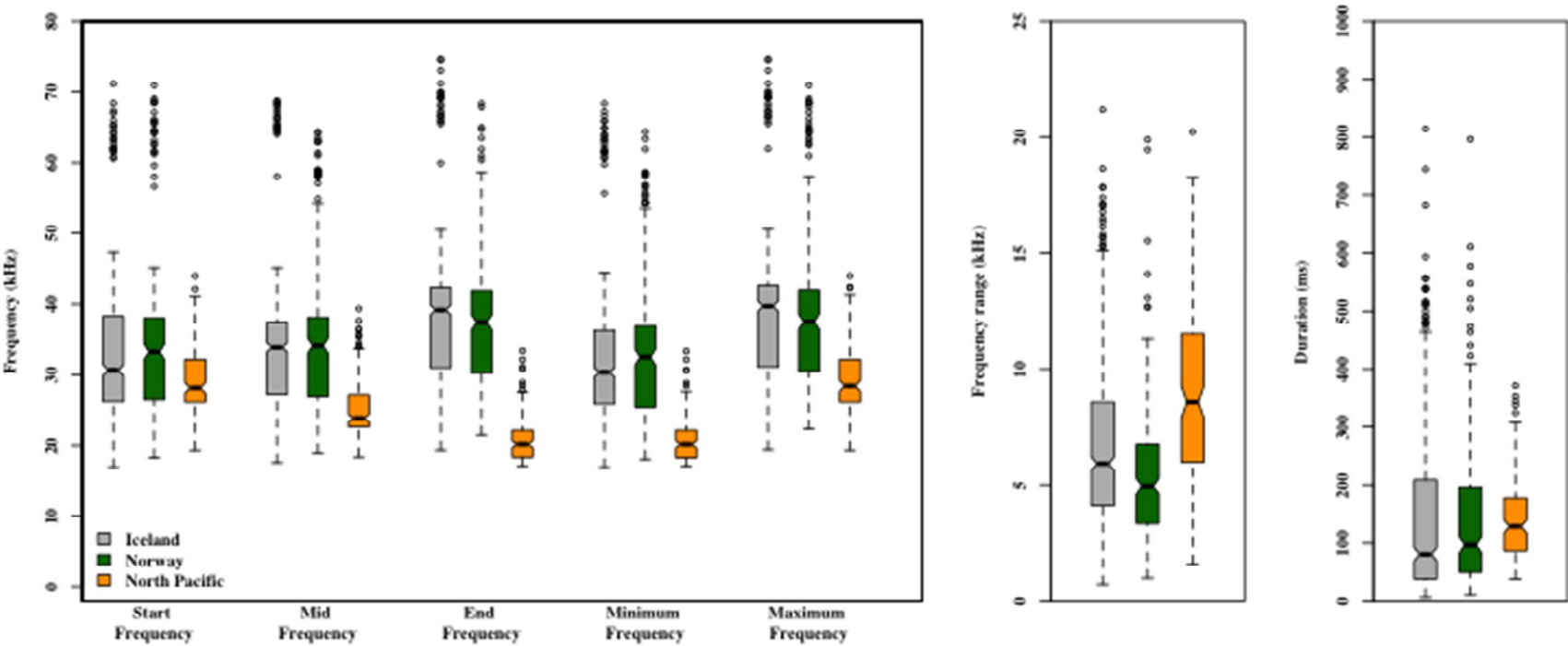
739

740 **Figure 2**



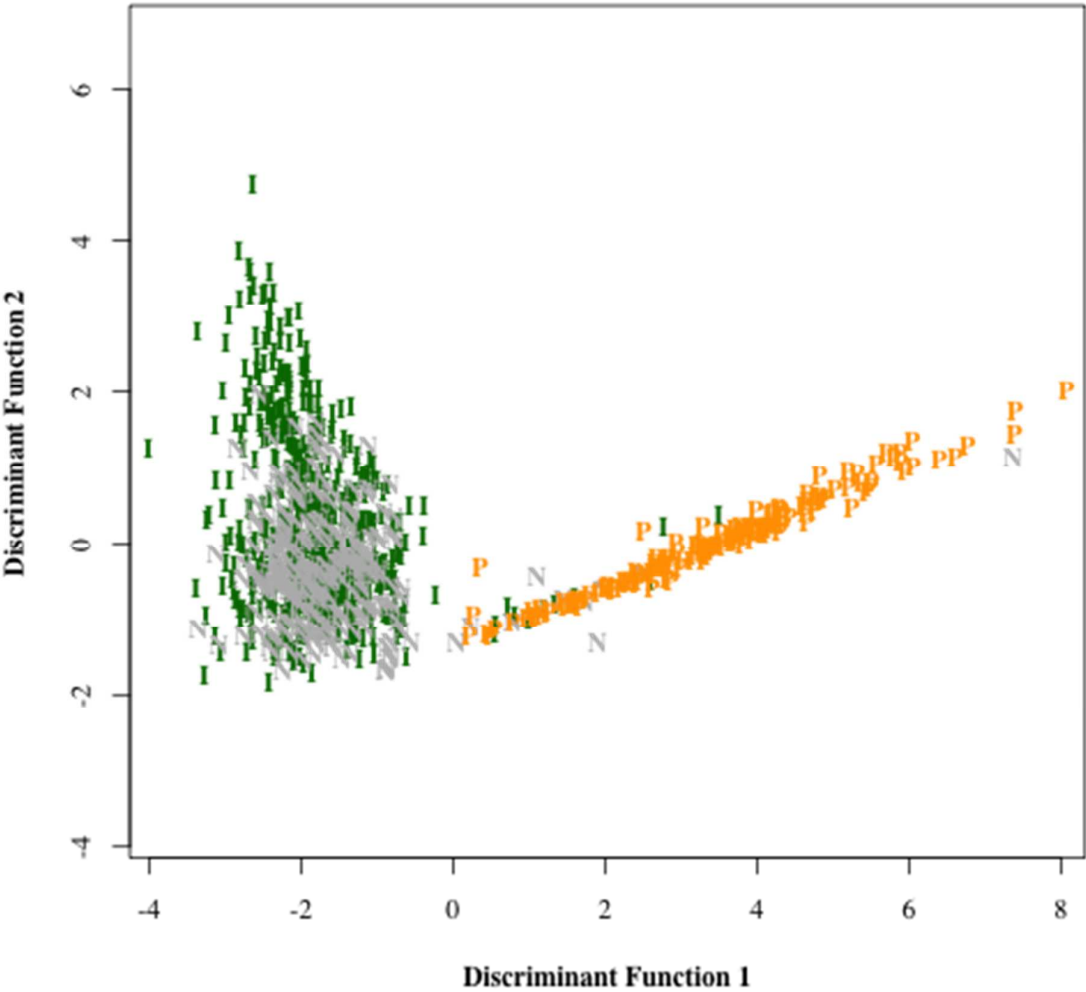
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742 **Figure 3**



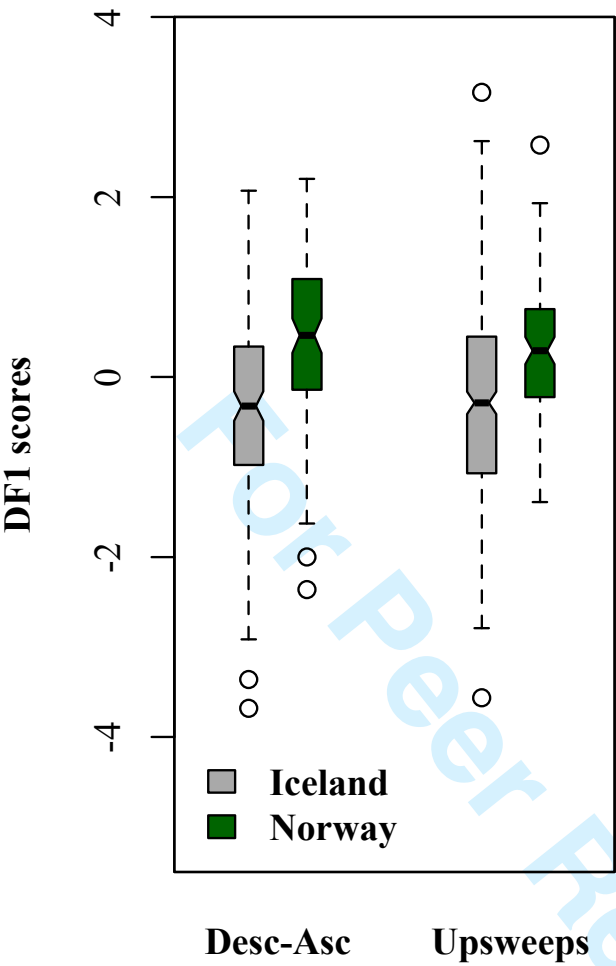
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744 **Figure 4**



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746 **Figure 5**



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